INTRODUCTION

Most coral reefs occur in oligotrophic waters with low dissolved nutrient concentrations and low standing stocks of phyto- and bacterioplankton. During passage over the coral reef, water features change and obtain a reef signature often characterized by depleted phytoplankton and bacterioplankton concentrations and enhanced dissolved nutrient concentrations (e.g. Ayukai 1993, Yahel et al. 1998, Gast et al. 1999, van Duyl et al. 2002). Growth rates of heterotrophic bacterioplankton in coral reef overlying waters are usually enhanced the closer the bacteria get to the reef and the coral bottom (Moriarty et al. 1985, Gast et al. 1999, van Duyl & Gast 2001). The highest growth rates have been reported for the coral surface microlayer (coral contact water) and coral mucus layer (Ducklow & Mitchell 1979, Paul et al. 1986, van Duyl & Gast 2001). Coral exudates are suitable food for the growth of heterotrophic bacterioplankton (Ferrier-Pagès et al. 2000). Besides a source of organic matter via benthic primary producers (corals and benthic algae), the reef bottom is a sink for particulate organic matter (e.g. plankton, detritus) and source of inorganic nutrients (Capone et al. 1992, Hatcher 1997, Rasheed et al. 2002). The benthic mineralization is fuelled by particulate organic matter settling on the bottom, by benthic microalgae (Heil et al. 2004), and by plankton trapped and mineralized by predominantly benthic heterotrophic suspension-feeding organisms, such as gorgonians, tunicates, and sponges (Buss & Jackson 1981, Bak et al. 1998, Pile 1999, Fabricius & Dommisse 2000, Wild et al. 2004). Hotspots for plankton depletion and mineralization on coral reefs are cryptic habitats, such as crevices.

ABSTRACT: Abundance, growth rates and nutrient limitation of the heterotrophic bacteria present in the reef water column and reef cavity water were measured on a fringing reef at Curáçao (Netherlands Antilles). Bacterial in situ growth rates were measured using dialysis bags. Nutrient limitation was measured using bioassays with different amendments of inorganic nutrients (nitrate, ammonium, phosphate) and dissolved organic carbon (glucose). Cell sizes were measured in the reef water column and over an intra-cavity gradient inside reef cavities, from the cavity center to the wall of the cavity. Bacterial abundance was lower in cavities, while growth rates were on average 3.6 times higher than in open reef water. The bacterial community in open reef water was limited in its growth by all nutrients, in contrast to cavity water, in which bacteria were limited by phosphate. Cell volumes decreased significantly from open reef water towards the cavity wall. Results suggest that conditions in cavities have a positive effect on bacterial growth. Growth limitation by N is alleviated through enhanced mineralization in cavities. Through rapid exchange with the ambient water, cavities enrich the overlying reef water with inorganic N.

KEY WORDS: DOC · DIP · DIN · Nutrient limitation · Growth rates · Heterotrophic bacteria · Cavities · Coral reef

Why is bacterioplankton growth in coral reef framework cavities enhanced?

S. R. Scheffers1, 2, 3, 4,*, R. P. M. Bak1, 3, F. C. van Duyl1

1Royal Netherlands Institute for Sea Research (NIOZ), PO Box 59, 1790 AB Den Burg, Texel, The Netherlands
2CARMABI Foundation, PO Box 2090, Willemstad, Curáçao, Netherlands Antilles
3University of Amsterdam, IBED, PO Box 94766, 1090 GT Amsterdam, The Netherlands
4Present address: University of Essen, Faculty of Biology and Geography, Universitätsstraße 5, 45141 Essen, Germany
and coral cavities (Richter et al. 2001, Scheffers et al. 2004). Despite high effluxes of ammonia and nitrate, inorganic nutrient concentrations usually remain low in the benthic boundary layer, due to efficient internal cycling and water exchange. Inorganic N and P concentrations in coral cavities are usually enhanced compared to concentrations on the coral reef bottom and in the overlying water (Gast et al. 1998, Richter et al. 2001). Bacterioplankton in the benthic boundary layer is confronted with these different gradients of nutrients, which can result in small-scale spatial variations in growth rates (van Duyl & Gast 2001).

It is now well established that nitrogen and, in particular, phosphorus may limit growth of heterotrophic bacteria in marine waters (Elser et al. 1995, Pomeroy et al. 1995, Thingstad & Rassoulzadegan 1995, Cotner et al. 1997, Thingstad et al. 1998, Sala et al. 2002). Bacteria acquire a significant fraction of their N and P demand via the uptake of inorganic forms of these elements (e.g. Kirchman 2000). Particularly in oligotrophic environments, inorganic nutrients are a potentially limiting factor for bacterial growth. The ability to take up inorganic nutrients places heterotrophic bacteria in direct competition with primary producers. Heterotrophic bacteria have P requirements that are typically 10 times higher than those of algae, whereas their N demand is not that different (Bratbak 1985, Vadstein et al. 1988, Lee & Fuhrman 1987). Therefore, bacteria can be an important competitor for P with algae and a sink for phosphorus. Bacterioplankton may become P-limited in environments where dissolved organic carbon (DOC) and N are available.

Metazoan grazing on bacteria and/or a poor ability to retain acquired P by bacterioplankton may alleviate bacterial P limitation (Vadstein et al. 2003). N and C limitation of bacterial growth appear to be less common, but also occur (Carlson et al. 2002, Sala et al. 2002). Bacterial growth in atoll lagoons appears to be primarily limited by inorganic N, but organic P and DOC limitation were also observed (Torréton et al. 2000). Carbon incorporation ultimately determines the N and P demand of growing bacteria (Carlson et al. 2002).

We examined heterotrophic bacterial standing stocks, bacterial growth rates, and bacterial nutrient requirements (through macronutrient and DOC amendments in batch cultures) in 2 different water types: reef cavity water and reef water column. The aim of our study was to investigate if the enhanced concentration of cavity inorganic nutrients contributes to the higher growth rate of reef cavity bacteria compared to reef water column bacteria.

**MATERIALS AND METHODS**

**Study area.** The framework cavities used in our study were located on the fringing reef of Curaçao, Netherlands Antilles (12° 12' N, 68° 56' W). We studied 10 different cavities at a depth of approximately 15 m (Fig. 1) on the reef slope at CARMABI Buoy Zero/Buoy One (van Duyl 1985). The cavities were scattered over 200 m along the coastline. The cavities were comparable in their biotic and physical characteristics to those described by Scheffers et al. (2003, 2004). The selected cavities had volumes of approximately 100 l, solid carbonate walls, and a sandy bottom. The main opening of the cavities was directed off the reef slope, and the hard surface area was covered (>60%) with sessile suspension feeders, such as sponges, polychaetes, bivalves, and tunicates. Cavity water is designated in this study as the body of water in a cavity. All cavity water samples were taken from the center of a cavity. Open reef water was sampled directly in front of a cavity opening, 1 m away from the substrate. Reef water was sampled before cavity water to avoid disturbance of the water properties. Sampling took place between 10:00 and 14:00 h. The samples were kept cold in the dark until processing. Growth rate experi-
Bacterial abundance. Samples (10 ml) taken to obtain heterotrophic bacterial concentration, were fixed with 0.2 µm filtered and buffered (sodium tetaborate, pH = 7.9) formaldehyde (final concentration: 0.7%). In the laboratory, the samples were stained with acridine orange and subsequently filtered (0.03 bar underpressure) onto 25 mm, 0.2 µm polycarbonate filters (Nuclepore) supported by a 0.45 µm cellulose acetate filter, within 24 h after sampling (Hobbie et al. 1977). The filters were mounted on slides in non-fluorescent immersion oil (Olympus). The microscopic slides were then stored in a freezer (–20°C). Bacteria were counted and sized with a Zeiss Axioskop epifluorescence microscope (1250× magnification). We used a grid of 36 × 36 µm, divided in 10 rows and 10 columns, for counting and sizing bacteria. At least 10 random grids per filter and at least 200 cells were counted.

Bacterial growth rates. Bacterial growth rate measurements were performed with dialysis bags (Spectrapor 4 dialysis membranes 12 000 to 14 000 MWCO) of 100 ml (Herndl 1993). Before use, the dialysis bags were pre-soaked in 0.2 µm filtered seawater for 1 h. Since all molecules <14 kDa can pass the membrane, a steady supply of in situ nutrients and DOC was secured. In a previous study (Ferrier-Pagès & Gattuso 1998), cellulose dialysis membranes proved to be sufficiently permeable to inorganic and organic nutrients (diffusive exchange of <1 h) and had no effect (negative or positive) on plankton growth rates. Water samples were taken in situ (inside and in front of the cavities) in 750 ml syringes and immediately brought to the boat at the surface, where the sampled water was filtered (0.03 bar underpressure). Duplicate dialysis bags were filled with 0.8 µm filtered cavity or reef water. Subsequently, dialysis bags were attached to frames and placed on the spot where the sample water originated (1 frame in the cavity and 1 just outside). Samples were taken with 10 ml syringes through a specially designed cap at T = 0, at T = 8 h, and at T = 24 h. Samples were then brought to the laboratory on ice and in the dark for bacterial counting. Experiments were conducted in 10 different cavities. Bacterial abundance at T = 0 and at T = 24 h, and the doubling times of bacteria in dialysis bags were compared between reef and cavity waters.

Cell volume gradients within the cavity were obtained using a Plexiglas ruler (25 to 50 cm length, dependent on cavity size) on which 3, equally dis-tanced, 10 ml syringes were attached. The plungers could be pulled out simultaneously. The actual positions of the syringes were in the center of the cavity, close to the cavity wall (5 cm away), and midway in-between. A reef water sample was taken before the cavity water samples. A total of 10 different cavities were sampled, 200 bacteria per sample were sized, totaling 8000 bacteria. The mean cell volume of heterotrophic bacteria was calculated by measuring the length (l) and width (w) of cells and assuming cells to be cylindrical, with a hemisphere of diameter w at each end:

\[
\text{Volume} = \frac{\pi}{4} (w^2) (l - w/3)
\]

Bioassays. To investigate nutrient limitation of bacterioplankton growth in reef and cavity waters, we conducted a series of controlled enrichment experiments with cavity and reef waters. We focused on the inorganic macronutrients N (nitrate and ammonium) and P (orthophosphate), and on DOC. Bioassay water was collected in situ with 750 ml syringes and brought to the laboratory, in darkness, for treatment. The cavity water samples from 10 different cavities were gently mixed and filtered (0.03 bar underpressure) over a pre-rinsed 0.8 µm polycarbonate (Nuclepore) filter to reduce grazing activity. Most of the heterotrophic bacteria (0.2 to 0.4 µm diameter) will pass this filter. After filtration, the water was poured into duplicate 1 l, acid- and distilled-water-washed, PC bottles. Reef water samples were treated in the same way. Each set of bottles (duplicates) received the amendments summarized in Table 1. We used a total of 36 bottles for the experiment (9 × 2 for reef water and 9 × 2 for cavity water).

The racks with bottles were covered with light impermeable foil and put back in the sea to be subjected

Table 1. Overview of the nutrient amendments

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Amendment</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>Dissolved inorganic nitrate: 10 µM NaNO₃</td>
</tr>
<tr>
<td>NH</td>
<td>Dissolved inorganic ammonium: 10 µM NH₄Cl</td>
</tr>
<tr>
<td>P</td>
<td>Dissolved inorganic phosphate: 1 µM Na₅P₃O₁₀ · 12H₂O</td>
</tr>
<tr>
<td>C</td>
<td>Dissolved organic carbon: 2.5 g l⁻¹ glucose (β-glucosemonohydrate), 3.1 g l⁻¹ pyruvate (CH₃COCOONa), 3.4 g l⁻¹ acetate (CH₃COONa), 1 ml of this mixture l⁻¹</td>
</tr>
<tr>
<td>N+P</td>
<td>10 µM NaNO₃ + 1 µM Na₅P₃O₁₀ · 12H₂O</td>
</tr>
<tr>
<td>N+C</td>
<td>10 µM NaNO₃ + 1 ml of a mixture of 2.5 g l⁻¹ glucose (β-glucosemonohydrate), 3.1 g l⁻¹ pyruvate (CH₃COCOONa), 3.4 g l⁻¹ acetate (CH₃COONa)</td>
</tr>
<tr>
<td>P+C</td>
<td>1 µM Na₅P₃O₁₀ · 12H₂O + 1 ml of a mixture of 2.5 g l⁻¹ glucose (β-glucosemonohydrate), 3.1 g l⁻¹ pyruvate (CH₃COCOONa), 3.4 g l⁻¹ acetate (CH₃COONa)</td>
</tr>
<tr>
<td>N+P+C</td>
<td>10 µM NaNO₃ + 1 µM Na₅P₃O₁₀ · 12H₂O + 1 ml of a mixture of 2.5 g l⁻¹ glucose (β-glucosemonohydrate), 3.1 g l⁻¹ pyruvate (CH₃COCOONa), 3.4 g l⁻¹ acetate (CH₃COONa)</td>
</tr>
<tr>
<td>CTRL</td>
<td>Control, unamended</td>
</tr>
</tbody>
</table>
to natural, in situ, temperature (ca. 27°C). Heterotrophic bacteria abundance was counted 8 h after the start of the experiment and subsequently every 8 h for 64 h. Data are presented as percent change in bacterial abundance relative to the control.

RESULTS

Bacterial abundance and growth rates

Bacterial abundance at $T=0$ in the dialysis bags was significantly lower in cavity water than in reef water ($t$-test for paired comparisons: $t = 5.4; p < 0.05$), indicating that most experiments with cavity water started off with a lower bacterial abundance (Fig. 2a). Bacterial abundance ranged from $2.9 \times 10^5$ to $5.9 \times 10^5$ cells ml$^{-1}$ in cavity water and from $4.4 \times 10^5$ to $6.2 \times 10^5$ cells ml$^{-1}$ in reef water. Cavity water bacterial abundance was more variable than reef water bacterial abundance. After 24 h of incubation in dialysis bags, bacterial abundance was significantly higher in cavities than in reef water ($t$-test for paired comparisons: $t = 4.1; p < 0.001$), indicating that most experiments with cavity water reached a higher bacterial abundance (in 9 of 10 experiments) after 24 h of incubation (Fig. 2b). Bacterial abundances at $T=24$ h ranged from $5.1 \times 10^5$ to $20.4 \times 10^5$ cells ml$^{-1}$ in cavity water and from $4.7 \times 10^5$ to $9.7 \times 10^5$ cells ml$^{-1}$ in reef water (Fig. 2b).

Doubling times of bacteria exposed to cavity water were significantly shorter (on average 3.6 times shorter) than doubling times of bacteria exposed to reef water ($t$-test for paired comparisons: $t = 7.9; p < 0.05$). The average doubling time of reef water bacteria was 82 h and ranged from 36 to 149 h; cavity water bacteria showed an average doubling time of 24 h and a range of from 10 to 44 h (Fig. 3). The doubling times of cavity and reef water bacteria exhibited high spatial and temporal variability, whereby the doubling time variability of cavities was lower than that of reef water.

Plotting doubling times of cavity bacteria versus doubling times of reef bacteria shows that the latter significantly (1-sample $t$-test: $R = 0.93; p < 0.0001$) increases with the increasing doubling times of cavity water bacteria (Fig. 4).

![Fig. 2. Comparison of bacterial abundance in reef water (RW) and cavity water (CW) (a) at the start ($T = 0$) and (b) end ($T = 24$) of the dialysis bag experiments ($n = 10$). Cavity numbers 1 to 5 at B0 and 6 to 10 at B1 (averages of 2 replicates with standard deviations). Note the difference in scales of the y-axes](image1)

![Fig. 3. Doubling times reef water bacteria (RW) and cavity water bacteria (CW) of 10 different cavities. Cavity numbers 1 to 5 at B0 and 6 to 10 at B1 (averages of 2 replicates with standard deviations)](image2)

![Fig. 4. Relation between the doubling times of cavity water bacteria (CW) and those of reef water bacteria (RW). Continuous line is a linear regression fit with $R = 0.77$ ($p < 0.001$, averages of 2 replicates with standard deviations). Dashed line represents the line along which bacterial doubling times in reef water are the same as those in cavity water](image3)
Bioassays

The bioassay data (Fig. 5) show that there were response differences to amendments of inorganic nutrients and DOC between incubations of cavity and reef water bacteria. Bacterial growth in reef water and cavity water samples was not different without amendment (control, Fig. 5a). The heterotrophic bacteria within the cavity water samples have a potentially higher specific growth rate and/or reach a higher level when +P (Fig. 5c), a combination of +N+P (Fig. 5h), and +N+P+C (Fig. 5i) are added, compared to the control. On the other hand, +N (Fig. 5b), +C (Fig. 5d), a combination of +N+C (Fig. 5e), and +NH (Fig. 5f) additions showed no clear stimulation in bacterial growth rates in cavity water compared to the control. Reef water bacteria show a positive response in growth to all amendments. Addition of +P (Fig. 5c), +P+C (Fig. 5g), and +N+P+C (Fig. 5i) had the same quantitative effect on bacterial growth in both water types. The amendment of +N and +NH (Fig. 5b & f) to reef water incubations showed the sole clear difference in response between reef water and cavity water bacteria. The addition to both water types of a combination of +N+P+C resulted in the highest abundance of heterotrophic bacteria (~9 × 10^6 cells ml⁻¹).

Cell volume and concentration gradient

Heterotrophic bacterial cell characteristics were different for different water types (Fig. 6), with cell abundance (Fig. 6a) and volume (Fig. 6b & c) decreasing into the cavity and towards the cavity wall, covered with cryptic suspension feeders. Reef water contained significantly larger cells than cavity water (1-sample t-test: t = 3.58; p < 0.001). Within the cavity, cell volumes significantly declined from the center of the cavity towards the cavity wall (1-sample t-test: t = 16.7; p < 0.001). The range in cell volume was largest in reef water and smallest in water close to the cavity wall. Mean bacterial cell volume in reef water was 0.085 µm³ (standard deviation: 0.042 µm³); in center cavity water, 0.075 µm³ (standard deviation: 0.028 µm³); mid-way towards the cavity wall, 0.059 µm³ (standard deviation: 0.023 µm³); and in wall cavity water, 0.037 µm³ (standard deviation: 0.015 µm³). Cell concentration showed a similar pattern with bacterial abundance, decreasing from reef water towards the cavity wall (Fig. 6a). Size-frequency distributions (Fig. 6c) show that only the smaller cell size classes (<0.08 µm³) of the bacterial community were still present in the cavity water close to the cavity wall and that size classes >0.08 µm³ were absent in the cavity wall samples. It is possible that the volume of individual cells was slightly under- or overestimated, depending on the degree of fluorescence of sized cells, but it is unlikely that the change in size of bacteria from the center towards the cavity wall is due to this purported artifact, which occurs at random.

DISCUSSION

We found large differences in bacterial abundance and doubling times between reef and cavity waters over small spatial scales (10 to 100 cm). The bacterial abundance in cavity water is significantly lower than that in reef water, which is in agreement with Gast et al. (1998), who found a similar pattern of lower concentrations in crevice water compared to reef water on the terrace between 6 and 8 m depth. The abundance of bacteria in reef water (ca. 5.3 × 10^5 bacteria ml⁻¹) is within the range generally found in coral reef waters (3.0 to 9.0 × 10^5 bacteria ml⁻¹; see e.g. Moriarty et al. 1985, Ayukai 1993, Bak et al. 1998, Gast et al. 1998). The consumption of heterotrophic bacteria by suspension-feeding cryptofauna with fluxes of up to 49 mg bacterial C m⁻² cavity surface area d⁻¹ (Scheffers et al. 2004), accounts for lower bacterial concentrations in cavity water.

Our data show a significant difference between the bacterial doubling times in cavity water and those in reef water, with cavity water bacteria doubling times being on average 3.6 times shorter than the doubling times in reef water. Doubling times in reef water (on average 82 h) were comparable to doubling times in reef water as measured by 3H-leucine incorporation at the same site by Gast et al. (1998), who found bacterial doubling times of ca. 80 h in reef water and slightly shorter doubling times (on average 75 h) in small crevices with volumes of <1 up to several liters. Our average cavity bacteria doubling times (ca. 24 h) were considerably shorter, but were in the same range as those found in the literature for coral reef waters. Ferrer-Pagès & Gattuso (1998) report doubling times of 6 to 24 h in Bora Bay, Japan; Moriarty et al. (1990) report doubling times of 6 to 8 h in the Bay of Carpenteria, Australia; and Landry et al. (1984) present doubling times of 13 to 20 h in Kaneho Bay, Hawaii. The initial bacterial abundance was significantly lower in cavity water than in reef water. This may have enhanced the growth rate in dialysis bags with cavity water (more food for rapid growth of bacteria). However, the total abundance of bacteria was, after 24 h of incubation, higher in cavity water bags than in reef water bags, indicating that there was more potential for formation of bacteria in cavity water than in reef water. This suggests that bacteria were supplied with extra nutrients (dissolved organic matter, inorganic
Fig. 5. Bioassay experiments. Graphs show the change in abundance of reef water (RW) and cavity water (CW) heterotrophic bacteria to different combinations of nutrient and/or dissolved organic carbon additions. Each line is corrected for the control and converted to the percentage compared to the control (+N, nitrate; +NH, ammonium; +P, phosphate; +C, dissolved organic carbon [averages of 2 replicates with standard deviations]).
Scheffers et al.: Enhanced bacterioplankton growth

...nutrients) in cavity water. So the bacterioplankton in dialysis bags are continuously exposed to this water and apparently grow faster than bacterioplankton in dialysis bags in open reef water, where inorganic nutrients released from cavities are rapidly diluted. In the real reef situation, bacteria, which are exposed to cavity water during the short residence in cavities, will grow somewhat faster than bacteria outside cavities. The ecological significance is that the specific growth rates of bacteria in the benthic boundary layer (BBL) water are slightly higher than those of bacteria further away from the bottom.

Cell volumes of bacterioplankton in cavity water were, on average, smaller than the cell volumes in reef water, due to selective removal of larger cells by suspension-feeding cryptofauna in coral cavities. Smaller cells do have a higher affinity for nutrients (Button & Robertson 2000). This may suggest that cavity water bacteria are limited to a lesser extent by very low natural nutrient concentrations than the larger reef water bacteria. This may have contributed to the shorter doubling times of bacterioplankton in cavity water than in reef water and to the higher abundance of bacteria in cavity water than in reef water after 24 h of incubation in dialysis bags. Taking all these aspects into account, we conclude that cavity water bears a more favorable signature for bacterial growth, in terms of higher nutrient availability, than reef water.

The positive relation between the doubling times of reef water bacteria and those of cavity water bacteria suggest that an exchange exists between these 2 water types affecting the bacterioplankton dynamics. It is widely recognized that water exchange is required for advecting food to the cryptic biota of coral cavities, which is critical for their survival (e.g. Buss & Jackson 1981). Water exchange rates of coral cavities have been reported to be quite high, with water residence time in cavities of 4 to 5 min on average (Richter et al. 2001, van Duyl et al. in press). This high water exchange implies that cavity water favorable (in terms of higher nutrient availability) to bacterial growth is rapidly mixed with reef water, giving reef water a distinct signature. Such an effect is diminished with increasing distance from the cavity and reef bottom (van Duyl & Gast 2001). The continuous release of water from the cavities, which is enriched in nutrients and depleted in bacterial abundance, contributes to the enhancement of bacterial growth in the overlying reef water and, thus, to the enhanced turnover of matter in reef ambient water. Nutrients are released by cryptic biota and efficiently sequestered by, e.g., bacterioplankton, which is subsequently removed again by cryptic suspension feeders. This process is important for the understanding of nutrient cycles in the reef ecosystem, particularly because cryptic habitats with

Fig. 6. Changes in (a) bacterial abundance, (b) biovolume, and (c) biovolume frequency distribution of the bacterial community from reef water towards cavity water close to the cavity wall [RW, reef water bacteria [n = 2000]; CW center, bacteria from water of the center of a cavity [n = 2000]; CW middle, bacteria from water mid-way between CW center and CW wall [n = 2000]; CW wall, bacteria close to the wall of a cavity [n = 2000]). Boxes in (b) show median ± 25 % ranges, whiskers indicate 10 and 90 %, circles are outliers outside the 10th and 90th percentiles. Note: bacterial volumes might deviate from real, absolute volumes. This is due to the methods applied, which may either over- or underestimate cell volumes depending on the degree of fluorescence of individual cells.

Cavities are net sources of inorganic nutrients on coral reefs (Richter et al. 2001, Scheffers et al. 2004). Inorganic nutrient concentrations (e.g. NO\textsubscript{x}, PO\textsubscript{4}) are usually higher in cavities and crevices than in reef overlying water (Gast et al. 1998, Richter et al. 2001). At the study site van Duyl & Gast (2001) and van Duyl et al. (in press) recorded DIN (dissolved inorganic nitrogen) concentrations that were, in most cases, higher in crevices (0.6 to 1.8 µM) than in the reef water in front of crevices (0.5 to 0.9 µM); the same pattern, but less predictable, was found for inorganic P (in crevices: 0.05 to 0.1 µM; in reef water: 0.02 to 0.04 µM). Regeneration of inorganic nutrients in cavities and crevices can be relatively rapid. Scheffers et al. (2004) have reported a net mean DIN release rate from cavities at the same study site in Curacao of 0.67 mmol m\textsuperscript{-2} cavity inner surface d\textsuperscript{-1}. For DIP (dissolved inorganic phosphorus), they did not establish a significant release from cavities. But net effluxes of DIP from cavities have been established (Richter et al. 2001, van Duyl et al. in press). These fluxes suggest that inorganic nutrient limitation may be alleviated in cavities, which could lead to the enhancement of bacterial growth in cavity water.

It is widely acknowledged that bacteria take up inorganic nutrients at low concentrations (Kirchman 2000). The ecological significance of this capacity has been recognized, and the mechanism is incorporated into models of nutrient and carbon flow in aquatic ecosystems (Bratbak & Thingstad 1985, Thingstad 1987). However, empirical evidence of spatial and temporal variability in nutrient limitation of bacterial growth in oligotrophic coral reef waters is still rare. Limitation of bacterial growth by nitrogen or phosphorus presumably occurs as a consequence of the utilization of substrates that have high C:N or C:P values (Goldman et al. 1987, Goldman & Dennett 1991, 2000). It has been suggested that labile organic carbon compounds may accumulate under these conditions and that these carbon sources are taken up rapidly and metabolized when the growth-limiting element becomes available (Cotner et al. 1997).

Our bioassay results show that inorganic N and P are potentially limiting factors in bacterial growth on reefs. Cavity water bacteria show a response of the community to enrichment with phosphate. The experiments with reef water bacteria indicate that the growth of the bacterial community is limited by all macronutrients (N, P, and organic C). Our bioassays show that cavity water bacteria are phosphate-limited, despite the fact that DIP concentration in crevices is usually enhanced compared to the overlying water (van Duyl & Gast 2001). A relatively high bacterial demand for P could explain the P limitation of cavity bacteria. Some studies describe inorganic N, instead of P, as the predominant growth-limiting factor of bacteria in coral reef waters (e.g. Torréton et al. 2000). We found that after P alleviation, inorganic N is the limiting factor for bacterial growth in cavity water bioassays. After P and N addition the bacterial abundance in cavity water bioassays exceeded the abundance of bacteria in reef water bioassays, suggesting that there was more potential for cell production in cavity water than in reef water. This was also found within the dialysis bag experiments. Whether this was due to more labile DOC in cavity water than in reef water cannot be concluded, since we did not assess bacterial biomass or bacterial respiration. Nitrogen is the only amendment in cavity water, which did not show an increase in bacterial abundance relative to the control. Neither nitrate nor ammonia amendments stimulated bacterial production, suggesting that N is not the primary limiting factor for bacterial growth in cavity water. Mineralization in cavities apparently provides the bacterioplankton with sufficient inorganic N to alleviate N limitation. This N supply is predominantly in the form of nitrate (Scheffers et al. 2004), which needs to be reduced before uptake. Nitrate and ammonia amendments rendered, however, comparable results in reef and cavity water bioassays.

Besides N and P, reef water bacteria appear to be predominantly DOC-limited. Bacterial abundance in reef water increased considerably after DOC amendments. Also, after addition of inorganic nutrients, reef water bacterial abundance shows an increase. The most likely explanation of this increase in bacterial abundance in water with amendments is that sub-communities within the reef water community react differently to different additions. Fonnès Flaten et al. (2003) suggest that in bioassay experiments with whole bacterial communities not all sub-communities react the same to nutrient and DOC amendments. Carlson & Ducklow (1996) showed DOC limitation of heterotrophic bacteria in the Sargasso Sea. Others have found that the addition of labile carbon together with inorganic nutrients can stimulate bacterial production more than the sole addition of labile carbon (Cotner et al. 1997 Shiah et al. 1998, Thingstad et al. 1998, Carlson et al. 2002), indicating that energy and nutrient limitation can co-exist. We found comparable results for bacterial abundance increase when N, P, and C were added at the same time in both crevice water and reef water. The results show a smaller in-
crease in bacterial abundance in reef water with only carbon amendment, indicating that once the energy requirement was alleviated a secondary limitation was evoked. Del Giorgio & Cole (1998) discussed the possibility that bacteria maximize their rate of carbon catabolism of available C, even when inorganic nutrients are not available. The potential advantage of this strategy of maintaining high electrochemical potentials across bacterial cell membranes is to support active transport functions and motility. Both of these functions would leave cells balanced to exploit transient increases in nutrients associated with the patchy nutrient distributions present in Curaçao reef water (Del Giorgio & Cole 1998, Gast et al. 1998).

All experiments were performed under grazed-reduced/excluded conditions. Nanograzers can recycle nutrients required by bacterioplankton (Strom 2000), but grazing of bacteria by nanoflagellates is insignificant, and nanoflagellate numbers in Curaçao reef water are low (Gast et al. 1999). If there were an effect of grazing by nanoflagellates on bacterial numbers and growth, it may only be discernable after >50 h of incubation (Wilcox & Fuhrman 1994). Our bioassay experiments were run for only 2.5 d, and the conclusions are based on processes occurring within the first 50 h. We assumed that the chemical form of these recycled nutrients and the N-containing products of sponge excretion are similar to those used in the experimental amendments. However, we cannot rule out the possibility that compounds (e.g. DON, DOP) other than NH₄, NO₃, and PO₄ in material released by grazers facilitate the utilization of otherwise unpalatable dissolved organic matter. Viral induced lysis of bacteria can have an effect on growth rates of susceptible bacteria; however, the literature shows that the overall bacterial density need not be affected (Middelboe et al. 2001). If a degree of bacteriophagy by marine viruses occurs, it would occur in all our experiments and tend to be evened out in our comparisons of the 2 different water types.

We conclude from the bioassay experiments that: (1) Bacterioplankton growth in coral cavity water is enhanced by alleviation of inorganic N limitation. (2) Cavity water bacteria communities are primarily P-limited. After alleviation of P limitation, N was the limiting nutrient. (3) The bacterial community in reef water is limited by the macronutrients N, P, and C.

Because the bacterial growth rates in reef water and cavity water are tightly coupled, we argue that the C/N/P-limited reef water bacterial community increases its specific growth rate during passage through coral cavities. Bacteria exposed to cavity water grow faster than bacteria exposed to reef water. This implies that bacteria, while present in regenerative spaces (cavities), grow faster than open reef water bacterioplankton.

In conclusion, our experiments show different responses of heterotrophic bacterial growth to characteristics of reef water column and reef cavity water. Specific bacterial growth is enhanced during its short residence in cavities through alleviation of N limitation. Our bioassays show that reef water bacteria are limited by dissolved organic carbon, inorganic nitrogen, and phosphorus, whereas bacterioplankton in cavity water primarily lack phosphorus. In view of the rapid exchange rate of cavities with the overlying reef water, our study supports the hypothesis that cavities play an important role in the enhancement of growth of the bacterioplankton that pass over coral reefs.

Acknowledgements. We wish to thank the staff of the CARMABI Foundation (Curaçao, Netherlands Antilles) and especially ‘Don’ Carlos Winterdaal and Brian Leysner for their cooperation. Gerard Nieuwland (NIOZ) and Arjen Kop (NIOZ) are thanked for their logistic support in the Netherlands and in Curaçao. Very special thanks go to Anja Schefers for her assistance in the field. The critical comments of the reviewers contributed to further improvement of the manuscript. This study was financed by the WOTRO (Netherlands Foundation for the Advancement of Tropical Research) Grant No. W84-439.

LITERATURE CITED


Scheffers et al.: Enhanced bacterioplankton growth

Editorial responsibility: Otto Kinne (Editor-in-Chief), Oldendorf/Luhe, Germany

Submitted: November 9, 2004; Accepted: May 10, 2005
Proofs received from author(s): August 1, 2005